## **LISTING OF CLAIMS**

1. (Currently Amended) A method of analyzing a table of data relating to gene expression that varies with time and/or with the changing of environmental conditions, for identifying groups of co-regulated and co-expressed genes, comprising:

defining a clustering criterion of the data of said table;

in function of said clustering criterion, identifying in sub-tables (CLUSTERS) groups of genes that satisfy said clustering criterion;

establishing pair combinations of said sub-tables;

calculating characteristic parameters of the data associated to the groups of genes of each pair combination;

generating for each <u>pair</u> combination a characteristic value in function of the characteristic parameters of the groups of genes by a decision algorithm based on soft computing techniques;

identifying the <u>pair</u> combinations <u>whose</u> the characteristic value <u>of which</u> is greater than a certain pre-established threshold as "Gene Networks" and discarding <u>pair</u> combinations of groups of genes <u>whose</u> the characteristic value <u>of which</u> is smaller than said threshold.

2. (Currently Amended) The method of claim 1, further comprising the operations of:

defining a number of logic filtering criteria of the data of said table;

for each logic filtering criterion, generating a corresponding filtered sub-table (FILTER) containing data of genes <u>having</u> the expression values of which satisfy said logic filtering criterion; and

establishing pair combinations of sub-tables generated by filtering the data of said table with said filtering criteria and by said clustering.

- 3. (Currently Amended) The method of claim 1, wherein said decision algorithm is a fuzzy logic algorithm <u>having</u> the antecedents and consequents of which are defined in function of said parameters.
- 4. (Currently Amended) The method of claim 1, wherein said parameters are chosen from the group comprising composed of numerical parameters tied to gene expression levels, parameters having a semantic biological meaning, and mixed parameters expressing at the same time a numerical relationship and a semantic meaning.

5. (Currently Amended) The method of claim  $\underline{4}$  1,

wherein said <u>numeric</u> parameters are chosen from a group consisting of:

absolute values of linear correlation coefficients among data associated to pairs of genes;

absolute values of quadratic correlation coefficients among data associated to pairs of genes;

percentage of genes of the combination the final value of gene expression of which is greater than the respective value of initial gene expression;

percentage of genes of the combination the final value of gene expression of which is smaller than the respective value of initial gene expression;

percentage of genes the values of gene expression of which have a same increasing or decreasing time evolution; and

percentage of genes that have a maximum value of gene expression in a same condition; and

wherein said parameters having a semantic biological meaning are chosen from a group consisting of:

percentage of genes that have ontologies in common; <u>and</u> percentage of genes that have functional domains in common.

6. (Original) The method of claim 1, further comprising discarding combinations among sub-tables constituted by a number of genes smaller than a certain pre-established number, considering only once the genes that are comprised in both combined sub-tables.

- 7. (Currently Amended) The method of claim 1, wherein said clustering criteria are based on algorithms chosen in <u>a</u> the set <u>comprising</u>: constituted by agglomerative hierarchic algorithms, non hierarchic Kmeans algorithms, hierarchic sequential Kmeans, non-hierarchic SOM and not exclusive Fuzzy Clustering.
- 8. (Currently Amended) The method of claim 5, comprising calculating correlation coefficients of all pairs of gene sequences of the combination; subdividing the interval from 0 to 1 in five sub-intervals of equal length and assigning to each of said sub-intervals a respective quantized value of correlation (vi);

calculating the percentage of correlation coefficients belonging to each sub-interval; defining for each combination an overall coefficient of linear correlation obtained as arithmetic mean of the quantized values associated to the sub-intervals containing a number of coefficient greater than 50%.

 (Currently Amended) The method of claim 5, comprising calculating coefficients of quadratic correlation of all pairs of gene sequences of a same combination;

defining for each combination a global coefficient of quadratic correlation obtained as <u>an</u> arithmetic mean of said correlation values.

## 10. (Original) The method of claim 5, comprising

calculating the percentage of sequence of the combination with a final value of gene expression greater than the initial value of gene expression;

defining a coefficient of global variation of the value of gene expression, comprised between 0 and 1, corresponding to said percentage.

## 11. (Original) The method of claim 5, comprising

calculating the percentage of sequences of the combination with an increasing time evolution;

defining a coefficient relative to the time evolution of the gene expression comprised between 0 and 1 corresponding to said percentage.

## 12. (Original) The method of claim 5, comprising

calculating the percentage of sequences of the combination with a value of gene expression greater than a pre-established threshold in correspondence of a same instant;

defining a coefficient of presence of maximum excursion of the level of gene expression in correspondence of the same instant, comprised between 0 and 1, corresponding to said percentage.

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13. (Currently Amended) An identification system of groups of co-expressed and coregulated genes, according to the method of claim 1, comprising:

a pre-processing sub-system input with data of a table relative to gene expressions variable with time and/or relating to different environmental conditions, the pre-processing sub-system generating sub-tables (CLUSTERS) of data in groups of genes that satisfy a pre-established clustering criterion;

a processing sub-system of data of said sub-tables (CLUSTERS), generating signals representing characteristic parameters of data associated to genes of a same combination of pairs of said sub-tables;

an intelligent sub-system input with said signals representative of characteristic parameters, generating for each combination a characteristic value in function of the characteristic parameters of the groups of genes and outputting data of groups of genes identified from combinations whose characteristic value is greater than a certain pre-established threshold as "Gene Networks."

14. (Original) The system of claim 13, wherein said intelligent sub-system is a neural fuzzy logic sub-system, trained off-line.